The production of fatty acid modifying enzyme (FAME) and lipase by various staphylococcal species

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Summary. Eighty-six strains encompassing 11 species of coagulase-negative staphylococci were examined for the production of fatty acid modifying enzyme (FAME) and lipase. Staphylococcus schleiferi and S. saprophyticus most closely resembled S. aureus in that 80% of the strains produced both enzymes. In contrast, no strains of S. lugdunensis and S. haemolyticus tested produced these enzymes. S. simulans was unusual in that eight of 10 strains produced FAME, but only one produced lipase. Among the other species the proportion of strains producing both enzymes ranged from 10 to 60%. Generally there was a strong correlation between FAME and lipase production.

Introduction

The multiplication and survival of Staphylococcus aureus within abscesses is controlled by the host through the production of two kinds of bactericidal lipids.1-3 Most strains of S. aureus produce an esterifying enzyme (fatty acid modifying enzyme; FAME) which can inactivate these bactericidal lipids and this enzyme appears to be important for the organism's survival in host tissues.4,5 FAME is inhibited by glycerides, and the host delivers large amounts of triglycerides into the core of abscesses which keeps this enzyme non-functional.2,6 However, if the organism can also elaborate lipase, this enzyme can relieve the inhibition. Therefore, to effectively negate the bactericidal activity of the lipids in lesions, the organism must be able to produce both FAME and lipase.

Currently, there are c. 28 known species of staphylococci other than S. aureus. Collectively, these are called coagulase-negative staphylococci (CNS), even though a few do produce coagulase, or clumping factor. Whereas it has been known for some time that some of these organisms can elaborate lipase,7,8 the production of FAME has not been evaluated.

Materials and methods

Staphylococcal cultures

Cultures of S. epidermidis, S. haemolyticus, S. warneri, S. lugdunensis, S. saprophyticus, S. simulans, S. schleiferi, S. caprae, S. kominis, S. cohnii and S. capitis were obtained either from Dr J. Fluerette at the National Staphylococcal Reference Center in Lyon, France, or from Dr L. Ayers at the University Hospital of The Ohio State University. Most strains had been isolated from closed-space lesions.

Evaluation of FAME production

Each strain was grown in 100-ml volumes of trypticase soy broth and was incubated at 37°C for 24 h with agitation. The culture supernates were divided into several portions and were kept frozen at -20°C until required.

For these studies, a simpler FAME assay was developed which measured the esterification of oleic acid to butanol and which quantified the ester by gas chromatography.4,5 A solution of oleic acid in HPLC-grade butanol (4 mg/ml) was prepared; 50 μl of this solution was placed in a lipid-clean glass screw-capped vial and to this was added 700 μl of 0.1 M sodium phosphate buffer (pH 6.0) and 250 μl of culture supernate. The mixture was incubated for 20 min in a water bath at 37°C with constant shaking. Each sample was extracted twice with 2-ml volumes of ethyl ether:methanol (6:1 v:v) and once with 2 ml of ethyl ether. The ether phases were pooled and dried under nitrogen. The extracted lipids were dissolved in 40 μl of hexane and 1-μl samples were injected into a gas chromatograph fitted with a 30-m carbowax 20 M capillary column (Ohio Valley, Marietta, OH, USA). Using helium as the carrier gas, the butyl ester eluted at 4.4 min and the free oleic acid at 13.0 min. The areas under the peaks were integrated and the percentages of free oleic acid and butyl ester were calculated. Culture supernates negative for FAME activity were re-tested to verify lack of enzyme production by the same procedure, except that the reaction mixture was incubated for 2 h instead of 20 min.

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One unit of FAME activity (butanol unit) was defined as the esterification of 10 nmol of oleic acid/min. Under the test conditions, this was equivalent to 28% esterification of the oleic acid during the 20-min incubation period.

**Evaluation of lipase activity**

A substrate suspension, similar to that used by Brunner and co-workers, was prepared by homogenising 2 ml of sodium taurocholate (15 mg/ml) with 5 ml of gum acacia 10% and 15 mg of triolein. To this was added 2 ml of 3 M NaCl, 1 ml of 0.075 M CaCl₂, and 4 ml of distilled H₂O. The mixture was adjusted to pH 8.0 while stirring and the final volume was adjusted to 15 ml with water.

To evaluate lipase production, 4 ml of culture supernate was added to 1 ml of substrate. The pH was again adjusted to 8.0 and the mixture was incubated in a water bath for 18 h at 37°C with constant agitation. After incubation, the lipids were extracted by the method of Bligh and Dyer and dried under nitrogen. The dried sample was dissolved in 240 µl of hexane and 20 µl of this was spotted on silica gel G-coated plates. The plates were developed in hexane:ethyl ether:acetic acid (80:20:1) and dried. Lipids were visualised by spraying with sulphuric acid:dichromate and charring at 180°C for 1 h. Lipid extracts of uninoculated medium with substrate as well as samples of pure triolein and oleic acid were included as controls. The procedure can detect the release of as little as 0.6% of the fatty acid in the substrate.

**Bactericidal activity**

The method for determining the sensitivity of a strain to the bactericidal lipids has been described in detail. In this study, oleic acid was used as a representative lipid because it is the predominant fatty acid in the free fatty acid pool produced in abscesses.

**Results**

Between five and 10 strains from each of 11 staphylococcal species were examined for FAME and lipase production (table I). Of these, *S. schleiferi* most closely resembled *S. aureus* in that 80% of the strains were strong producers of both FAME and lipase. Five of six *S. cohnii* strains and seven of nine *S. saprophyticus* strains also produced both enzymes. In contrast, no strains of *S. lugdunensis* and *S. haemolyticus* tested produced FAME or lipase. *S. simulans* was unusual in that, whereas eight of 10 strains were strong FAME producers, only one of these strains could elaborate lipase. Among the other strains, enzyme production was less consistent.

Although only 19 strains were tested for sensitivity to oleic acid, most were found to be sensitive (table II) and a few, such as the two *S. lugdunensis* strains, were inordinately sensitive to the fatty acid.

**Discussion**

Although the system of bactericidal lipids plays an important role in controlling the growth and survival of *S. aureus* in focal lesions, it is not known whether this host defence mechanism is brought to bear on other species of staphylococci. In the case of *S. aureus*, two enzymes assist the organism in circumventing these lipids. The first is FAME, an esterifying...
enzyme that can inactivate both the free fatty acid pool and the monoglycerides. \(^5\) However, FAME is strongly inhibited by triglycerides and the host consigns large amounts of triglycerides to the lesion. The second enzyme, lipase, can relieve this inhibition and can also degrade the bactericidal monoglyceride directly. Therefore, to obtain the greatest benefit, the organism should produce both enzymes simultaneously. Whereas about 99.5% of \(S.\) aureus strains produce lipase,\(^6\) only 80% produce FAME,\(^6\) yet it is only these strains that are able to invade the tissues.\(^4\)

In the present study, c. 80% of \(S.\) schleiferi, \(S.\) cohnii, and \(S.\) saprophyticus strains produced both FAME and lipase, but none of the \(S.\) haemolyticus and \(S.\) lugdunensis strains produced either enzyme. Among the other species, the proportion of strains producing both enzymes ranged from 10 to 60%.

Generally, there appeared to be a strong correlation between FAME and lipase production (table I). Of a total of 86 strains examined, 33 (38%) produced both enzymes, and 35 (41%) produced neither. Only 18 strains (21%) produced one enzyme but not the other (\( \chi^2 \) analysis: \( p < 0.001 \)). This kind of relationship might be expected if the host utilised the defence mechanism against certain species, but not against others. Since production of a single enzyme would be of little value to a strain in negating the effect of bactericidal lipids, selective pressures would favour strains producing both enzymes in those cases where the defence mechanism was being utilised, but no such pressures would come into play in instances where the mechanism was not employed.

Of the 19 strains evaluated for their sensitivity to oleic acid (table II), 11 strains produced both FAME and lipase. The average level of sensitivity to the fatty acid for this group was 1030 LD50/mg (median, 910 LD50/mg). In contrast, six strains produced neither enzyme and in this group the average level of sensitivity was much greater, 5200 LD50/mg (median, 5375 LD50/mg).

Although our findings are compatible with the concept that at least some species of CNS may produce FAME and lipase for defensive purposes, further studies are obviously necessary to determine the validity of this view. Many infections due to CNS are associated with the presence of various types of foreign bodies, particularly those made of certain plastics. In addition to determining whether the host utilises, or attempts to utilise, the system of bactericidal lipids in response to infection with CNS, the effect of plastics on the mobilisation process or on the efficacy of the lipids must also be considered.

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References